Subpart B—Submission of Information

§805.10 Submission of information by physicians and providers.

A physician or a provider of services that requests or receives payment from Medicare for the implantation, removal, or replacement of a permanent cardiac pacemaker device or pacemaker lead shall submit the following information on a specified form to HCFA for inclusion in the pacemaker registry provided for by FDA under §805.1:

- (a) Provider number.
- (b) Patient's health insurance claim number (HICN).
 - (c) Patient's name.
 - (d) Date of the procedure.
- (e) Identification number (used by PRO's) and name of the physician who ordered the procedure.
- (f) Identification number (used by PRO's) and name of the operating physician
- (g) For each device (pulse generator, atrial lead, ventricular lead) implanted during the procedure about which the report is being made: the name of the manufacturer, model number, serial number, and the warranty expiration date.
- (h) For each device (pulse generator, atrial lead, ventricular lead) removed or replaced during the procedure about which the report is being made: the name of the manufacturer; model number; serial number; the warranty expiration date, if known; the date the device was initially implanted, if known; whether a device that was replaced was left in the body; if the device was not left in the body, whether it was returned to the manufacturer.

(Information collection requirements approved by the Office of Management and Budget under control number 0910–0234)

§805.20 How to submit information.

Information shall be submitted to the registry in the form and manner required under general instructions of the Medicare program (see 42 CFR 409.19(a) and 410.64(a)).

$\S 805.25$ Confidentiality.

(a) FDA and HCFA will keep confidential, and will not reveal to the

public, any specific information that identifies by name a recipient of any pacemaker device or lead or that would otherwise identify a specific recipient.

(b) Public disclosure of all other information under this part will be governed by the Freedom of Information Act (5 U.S.C. 552), the Privacy Act of 1974 (5 U.S.C. 552a), the Department of Health and Human Services' public information regulations (45 CFR part 5), FDA's public information regulations (21 CFR part 20), and HCFA's public information regulations (subpart B of 42 CFR part 401).

PART 807—ESTABLISHMENT REG-ISTRATION AND DEVICE LISTING FOR MANUFACTURERS AND DIS-TRIBUTORS OF DEVICES

Subpart A—General Provisions

Sec.

807.3 Definitions.

Subpart B—Procedures for Domestic Device Establishments

- 807.20 Who must register and submit a device list.
- 807.21 Times for establishment registration and device listing.
- 807.22 How and where to register establishments and list devices.
- 807.25 Information required or requested for establishment registration and device listing.
- 807.26 Amendments to establishment registration.
- 807.30 Updating device listing information.
- 807.31 Additional listing information.
- 807.35 Notification of registrant.
- 807.37 Inspection of establishment registration and device listings.
- 807.39 Misbranding by reference to establishment registration or to registration number.

Subpart C—Registration Procedures for Foreign Device Establishments

807.40 Establishment registration and device listing for U.S. agents of foreign manufacturers of devices.

Subpart D—Exemptions

807.65 Exemptions for device establishments.